## Pine Chemicals Association, Inc.

1117 Perimeter Center West Suite 500E Atlanta, GA 30338

July 23, 2001

Administrator USEPA P.O. Box 1473 Merrifield, VA 22116

Re: HPV Test Plan and Robust Summaries for Rosins and Rosin Salts

Dear Ms Whitman;

On behalf of the member companies of the Pine Chemicals Association's High Production Volume Chemical Task Force, I am pleased to submit the Test Plan and Robust Summaries for the chemical category designated "Rosins and Rosin Salts".

The submission includes one electronic copy in pdf. format, and a hard copy which will be mailed to EPA Headquarters. The registration number for our Consortium is

Please note from the above letter head that the headquarters of the Pine Chemicals-Association has moved to a new location with new contact information.

Should you have any questions concerning our submission please feel free to contact me at (770) 399-3112 or at wjones@pinechemicals.org.

Sincerely,

Walter L. Jones President & COO

v: 770.399.3112 f: 770.399.3115 2001 JUL 27 AN 9:

OPPT NCIC



Walter Jones <wjones@pinechemicals.org> on 09/10/2001 10:06:16 AM

Please respond to wjones@pinechemicals.org

To: NCIC OPPT/DC/USEPA/US@EPA, Rik Chem/DC/USEPA/US@EPA

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Subject: RESUBMISSION - TEST PLAN & ROBUST SUMMARIES - ROSIN AND ROSIN SALTS

Attached is a resubmission of the Test Plan and Robust Summaries for Rosin and Rosin Salts by the Pine Chemicals Association's HPV Task Force. The original submission of this test plan was made on July 26, 2001.

Information contained in the original submission concerning biodegradation data and future testing of Hydrogenated Rosin (CAS No. 65997-06-0) was incorrect.

Walter L. Jones, President & COO

Pine Chemicals Association, Inc.

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Rosin and Rosin Salt Test Plan.pdf

in PH12:3

# AR201-13134A

# HIGH PRODUCTION VOLUME (HPV) CHEMICAL CHALLENGE PROGRAM

## **TEST PLAN**

for

## ROSINS AND ROSIN SALTS

CAS No. 8050-09-7 CAS No. 65997-06-0 CAS No. 68425-08-I CAS No. 61790-50-9 CAS No. 61790-51-O CAS No. 68783-82-4

2001 JUL 27 AM 9: 46

Submitted to the US EPA

Ву

The Pine Chemicals Association, Inc.
HPV Task Force
Consortium Registration

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#### **Test Plan for Rosins and Rosin Salts**

#### Summary

The Pine Chemicals Association, Inc. (PCA) is sponsoring 36 HPV chemicals. This Test Plan addresses the following six chemicals, known collectively as Rosins and Rosin Salts:

CAS 8050-09-7, Rosin
CAS 65997-06-0, Rosin, hydrogenated
CAS 68425-08-1, Rosin, distillation overheads
CAS 68783-82-4, Rosin, low boiling fraction
CAS 61790-50-9, Rosin, potassium salt
CAS 61790-51-0, Rosin, sodium salt

All of the members of this group of substances are closely related to rosin, which is a naturally occurring substance found in trees, predominantly pine trees. Rosin is composed primarily of resin acids, a class of tricyclic carboxylic acids, but also contains minor amounts of dimerized rosin and unsaponifiable matter. Because of the complex nature of their composition, rosin and the rest of the compounds in this group are considered to be "Class 2" substances.

Rosin is an important commercial material and has been used for centuries. Industrially, there are three different types of rosin: gum, wood and tall oil rosin, with the name indicating the way in which the rosin is extracted from the tree. In the United States, tall oil rosin is by far the most commercially important variety. Rosin (CAS# 8050-09-7) also now includes CAS# 8052-10-6, rosin, tall oil, which was used in the 1990 IUR reporting on which the HPV program was based, pursuant to EPA's letter to PCA of March 13, 1992.

Rosin is used primarily as the raw material for the production of rosin derivatives, which go into the production of a wide variety of industrial products. For example, the largest use of rosin is in the production of derivatives for printing inks, adhesives, and coatings. Rosin salts are widely used in paper sizing. Other members of the group are used to produce soaps and detergents or to impart enhanced stability to specialty rosin-based adhesives.

Existing data on rosin and hydrogenated rosin already exist. Both rosin and hydrogenated rosin are non-toxic in acute toxicity tests in multiple species. Existing data from repeat-dose studies, including long-term carcinogenicity studies on both of these compounds, show low toxicity and no potential carcinogenic or reproductive effects.

Where applicable, PCA will conduct physical/chemical property and environmental fate testing on five of the substances in the group. (Two of the substances, rosin distillation overheads and rosin low boiling fraction, are nearly identical so that physical/chemical

testing will only be conducted on one of them.) PCA has elected to treat this group of chemicals as a category for purposes of the HPV program. Therefore, a representative of the category will be used for ecotoxicity and developmental toxicity testing (the additional tests needed to complete the SIDS endpoints).

Rosin (CAS# 8050-09-7) has been selected as the representative substance in this category for testing for the additional SIDS data. Rosin represents by far the greatest production volume, with almost four times more rosin manufactured than all other substances in this category combined. In addition, rosin is the raw material from which all the other category members are derived. Consequently, test results obtained on rosin will be most representative of the category.

PCA has reviewed existing data on these compounds. Available data demonstrate that both rosin and hydrogenated rosin have low toxicity following acute oral exposure. A number of repeat dose and long-term carcinogenicity studies also show low toxicity and no evidence of carcinogenicity for both rosin and hydrogenated rosin. Because neither rosin nor hydrogenated rosin were carcinogenic in long-term feeding studies, it is reasonable to conclude that neither of these substances is genotoxic. The sub-chronic studies demonstrating a lack of any toxicity to the reproductive organs fulfill the SIDS endpoint for reproductive toxicity. Because there are no data on developmental toxicity, rosin will be tested to fulfill this endpoint.

A brief summary of the available data for the substances in this category, and the anticipated additional testing, is described below in Table 1.

Table 1

Matrix of Available Adequate Data and Proposed Testing
On Rosins and Rosin Salts\*

					Req	uired SII	OS Endpo	oints							
Chemical and CAS #	Partition Coef.	Water Sol.	Biodeg.	Acute Fish	Acute Daph.	Acute Algae	Acute oral	Repeat Dose	In vitro genotox (bact.)	In vitro genotox (non- bact)	Repro/ Develop				
8050-09-7 Rosin	Test	Test	Adeq.	Test	Test	Test	Adeq.	Adeq.	Adeq.	Adeq.	Adeq. Repro/ Test Develop.				
65997-06-0, Rosin, hydrogenated	Test	Test	Test	С	С	С	Adeq.	Adeq.	Adeq.	Adeq.	С				
68425-08-1, Rosin, distillation overheads	Test	Test	Test	С	С	С	С	С	С	С	С				
68783-82-4 Rosin, low boiling fraction	No test	No test	No test	С	С	С	С	С	С	С	С				
61790-50-9, Rosin, potassium salt	Test	Test	Test	С	С	С	С	С	С	С	С				
61790-51-0, Rosin, sodium salt	Test	Test	Adeq.	С	С	С	С	С	С	С	С				

Adeq. Indicates adequate existing data

**Test** Indicates proposed testing

No test See test plan; essentially identical to rosin, distillation overheads.

C Indicates category read-down from existing or proposed test data on rosin.

\* No testing will be conducted for melting point, boiling point, vapor pressure, hydrolysis, photodegradation and transport and distribution between environmental compartments as explained in the test plan.

#### **Physical/Chemical Properties**

Physical and chemical properties will be determined when appropriate. However, many of the physical and chemical properties are either inappropriate or cannot be measured for these compounds:

 Melting points will not be determined because these substances are complex mixtures and either will not give a sharp melting point when heated or will decompose before they melt.

- Boiling points cannot be determined because these substances are complex mixtures and will decompose before they boil.
- <u>Vapor pressure</u> of these chemicals under ambient conditions is essentially zero and experimental measurement is not possible.
- Water solubility of five of the compounds in this category will be determined.
- <u>Partition coefficients</u> will be tested for five of the substances for which data do not already exist. The partition coefficient testing likely will yield a range of values representing the various components, rather than a single value representing the mixture.

#### **Environmental Fate**

With respect to the SIDS environmental fate endpoints:

- <u>Biodegradation</u> data will be generated for three of the compounds for which data are not already available.
- <u>Hydrolysis</u> in water will not be determined for any of the compounds in this
  category because the members of this category lack a functional group that
  would be susceptible to hydrolysis.
- <u>Photodegradation</u> is not relevant, since the vapor pressure of these compounds is essentially zero and they could not enter the atmosphere.
- <u>Transport and distribution between environmental compartments</u> will not be determined due to the inability to provide usable inputs to the required model.

#### **Ecotoxicity**

 Existing ecotoxicity data are not reliable due to inconsistencies in, or artificial methods of, sample preparation. Consequently, using rosin, <u>acute toxicity to fish</u>, <u>daphnia and algae</u> will be retested under conditions that maximize solubility, but reduce exposure to insoluble fractions, which may cause nonspecific toxicological effects.

### **Mammalian Toxicity**

- For the SIDS human health endpoints, there are adequate data on <u>acute toxicity</u>, <u>repeat dose toxicity</u>, <u>and reproductive effects</u> for both rosin and hydrogenated rosin. Rosin and hydrogenated rosin have been shown to be non-toxic in these tests.
- The availability of two-year feeding studies on rosin and hydrogenated rosin showing a lack of carcinogenicity obviates the need for in vitro genotoxicity testing.
- A <u>developmental toxicity</u> study on rosin will be undertaken to fulfill this SIDS endpoint.

The Pine Chemicals Association, Inc. HPV Task Force includes the following companies:

Akzo Nobel Resins

Akzo Nobel - Eka Chemicals Incorporated

Arizona Chemical Company

Asphalt Emulsion Manufacturers Association

**Boise Cascade Corporation** 

Cognis Corporation

Eastman Chemical Co. (including the former Hercules Inc. Resins Division)

Georgia-Pacific Resins Inc.

ICI Americas (including the former Uniqema)

Inland Paperboard & Packaging, Inc.

International Paper Co. (including the former Champion International Corporation)

Koch Materials Co.

McConnaughay Technologies, Inc.

Mead Corp.

Packaging Corporation of America

Plasmine Technology, Inc.

Raisio Chemicals

Rayonier

**Riverwood International** 

Smurfit – Stone Container Corporation

Westvaco

Weyerhaeuser Co.

The Task Force will be filing multiple test plans covering various chemicals. Not all members of the Task Force produce the substances covered by this test plan.

## I. Description of Rosins and Rosin Salts

The Pine Chemicals Association, Inc. (PCA) is sponsoring six HPV chemicals known collectively as Rosins and Rosin Salts. This category of chemicals consists of the following:

8050-09-7, Rosin 65997-06-0, Rosin, hydrogenated 68425-08-1, Rosin, distillation overheads 68783-82-4, Rosin, low boiling fraction 61790-50-9, Rosin, potassium salt 61790-51-0, Rosin, sodium salt

All of the members of this group are derived from rosin, which is a naturally occurring substance found in trees, predominantly pine trees. Rosin is composed primarily of resin acids, a class of tricyclic carboxylic acids, but also contains minor amounts of dimerized rosin and unsaponifiable matter. As complex mixtures, rosin and its derivatives are all considered as Class 2 substances.

Rosin is an important commercial material and has been used for centuries. Industrially, there are three different types of rosin: gum, wood and tall oil rosin, with the name indicating the way in which rosin is extracted from the tree. Gum rosin is obtained by slashing the tree and collecting the gummy exudates (oleoresin). This exudate consists of a mixture of rosin and turpentine and the rosin is recovered by distilling away the turpentine. Wood rosin is obtained by the solvent extraction of pine wood. Tall oil rosin is obtained by the distillation of tall oil, a by-product from the alkaline pulping of pine wood. In the United States, tall oil rosin is by far the most commercially important form of rosin.

The three rosins are chemically very similar. They all contain the same resin acids but the ratio of the acids is different. The difference arises because some of the resin acids are thermally unstable and isomerize to other resin acids during the production process. In 1991, the Pine Chemicals Association (then the Pulp Chemicals Association) proposed to the EPA that there should only be one CAS registry number to describe all three types of rosin. In a letter dated March 13, 1992, the Inventory Section of the EPA agreed with this request stating, "... rosin, CASRN 8050-09-7 will cover all types of rosin, irrespective of their method of production." Subsequently, the industry and the EPA have only used one CAS number for rosin. Thus, CAS# 8050-09-7 now also includes substances formerly reported as CAS# 8052-10-6.

#### A. Composition

Each species of pine tree has a somewhat different mix of resin acids. Even within a species, the mix of resin acids may be influenced by the climate and local terrain. However, all the members of this group are derived from rosin. "Hydrogenated rosin," as the name implies, is made by the catalytic hydrogenation of rosin. "Rosin, distillation

overheads" is formed as a by-product when rosin is processed at high temperatures and is made up primarily of resin acids and decarboxylated resin acids. Rosin, low boiling fraction is essentially identical to rosin, distillation overheads. The sodium and potassium salts are simply rosin that has been reacted with the appropriate base.

The general characteristics and composition of each of the substances in this category are addressed below.

#### 1. Rosin (CAS# 8050-09-7)

Rosin is a pale yellow, glass-like solid. The description of rosin listed in Appendix A of the TSCA Inventory is "A complex combination derived from wood, especially pine wood. Composed primarily of resin acids and modified resin acids such as dimers and decarboxylated resin acids. Includes rosin stabilized by catalytic disproportionation."

The composition of a typical tall oil rosin is shown in Table 2. As is evident, it consists of several major components and some minor ones. Rosin also contains trace quantities of numerous other components. Due to its complex composition, rosin is classified as a Class 2 substance. The structures of some of the more important resin acids found in rosin are shown in Figure 1.

Commercially, rosin is rarely categorized by its composition. Rather, it is usually specified by its softening point, acid value, and color (Zinkel and Russell 1989). In fact, the Naval Stores Act of 1923 (7 USC §§ 91-99), as amended in 1951, and regulations promulgated thereafter by USDA list only color as a specification for rosin.

Table 2

Composition of a Typical Tall Oil Rosin

Resin acid	Composition
Pimaric Sandarcopimaric Communic Palustric Isopimaric Abietic	4% 4% 1% 8% 11% 38%
Dehydroabietic Neoabietic Other compounds <sup>a</sup>	18% 3% 12%

a: other resin acids, high boiling fatty acids and unsaponifiable matter.

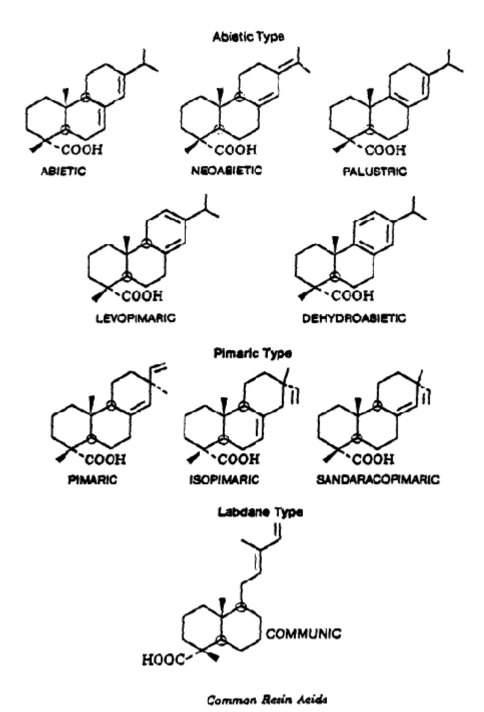


Figure 1. Representative resin acids found in rosin and its derivatives.

#### 2. Rosin, hydrogenated (CAS# 65997-06-0)

The composition of hydrogenated rosin is similar to rosin except that some of the double bonds in the resin acids have been removed. The conjugated double bonds in resin acids such as abietic acid are prone to oxidation and so catalytic hydrogenation is used to stabilize the molecule. The resulting product retains its color and other oxidation sensitive properties better than unmodified rosin. Like rosin, hydrogenated rosin is characterized primarily by its color, softening point stability and acid value rather than its chemical composition.

# 3. Rosin, Distillation Overheads (CAS# 68425-08-1) and Rosin, Low Boiling Fraction (CAS# 68783-82-4)

Rosin, distillation overheads, is one of several substances listed in the inventory that describe the product obtained when rosin is heated to the temperature at which it degrades. Another, virtually identical, substance is rosin, low boiling fraction. Descriptions of both these substances are listed in Appendix A of the TSCA Inventory and the descriptions of each are essentially the same. Rosin, distillation overheads, is described as "the low boiling fraction obtained by the distillation of rosin. Contains decarboxylated rosin, decarboxylated resin acids, resin acids, terpenes and hydrocarbons derived from decarboxylated fatty acids." Rosin, low boiling fraction is described as "A complex combination obtained by the distillation of rosin. This low boiling fraction consists primarily of decarboxylated rosin, resin acids, decarboxylated resin acids, terpenes and hydrocarbons derived from decarboxylated fatty acids."

Based on this similarity, testing for chemical/physical properties will only be conducted on rosin, distillation overheads and no testing will be done on the rosin, low boiling fraction. As these substances are by-products of other processes the composition can vary widely as shown in Table 3.

Table 3

Composition of a Typical Rosin, Distillation Overheads

Fatty acids	1-5%
Rosin acids	30-60%
Hydrocarbons <sup>a</sup>	20-40%
Rosin aldehydes	10-20%
Rosin alcohols	5-10%
Rosin esters	1-10%

a: terpenic and from decarboxylated rosin acids and fatty acids

## 4. Rosin, Potassium Salt (CAS# 61790-50-9) and Sodium Salt (CAS# 61790-51-0)

The rosin potassium and sodium salts are merely the simple alkali metal salts of unmodified rosin and are made by treating rosin with the appropriate base. As these substances are salts of a strong base and a weak acid they are alkaline with the pH depending on the concentration.

#### B. Commercial Uses of Rosins and Rosin Salts

Rosin is by far the most important member of this category from a commercial standpoint, with almost four times the volume of production as the other members of the group combined. The main use of rosin is in the production of derivatives or chemical intermediates that find a wide variety of industrial applications. The largest single application for rosin derivatives is in the production of printing inks, followed by adhesive, chewing gum, and coatings.

<u>Salts of rosin</u> are widely used in the paper and the soap and detergents industries. The sodium salts of rosin are used in paper sizing chemicals to give the finished product a better surface finish and water resistance. Potassium salts of rosin are used in the production of various soaps and detergents.

Rosin, hydrogenated is used in specialty adhesive applications where product stability and color are important. It is useful for these applications because it does not oxidize as readily as rosin.

Rosin, distillation overheads and rosin, low boiling fraction find application in the production of rosin derivatives for the end use applications described above, or if the quality of the substances is undesirable, they may be consumed for their fuel value.

## C. Complexity of Analytical Methodology

All of the substances in this category are Class 2 substances. This, combined with the fact that rosin is essentially insoluble in water and decomposes on heating at high temperature, creates a variety of analytical challenges. Gas chromatography of methylated derivatives is the accepted method for the analysis of the members of this category. PCA has verified the reliability of the standard analytical methods at concentrations lower than the 10 ppm limit of solubility for these substances. Based on the method validation work to date, it appears that the analytical procedures will be adequate for the proposed testing.

## II. Rationale for Selection of Representative Compound for Testing

Rosin (CAS# 8050-09-7) has been selected as the representative substance in this category for testing for the applicable SIDS ecotoxicity and developmental toxicity tests, as shown in Table 4 (identical to Table 1). As further indicated in Table 4, pertinent

physical/chemical properties and environmental fate endpoints will be determined for all five members of this category where data are not already available.

Table 4
Matrix of Available Adequate Data and Proposed Testing
On Rosin and Rosin Salts\*

		Required SIDS Endpoints									
Chemical and CAS #	Partition Coef.	Water Sol.	Biodeg.	Acute Fish	Acute Daph.	Acute Algae	Acute oral	Repeat Dose	In vitro genotox (bact.)	In vitro genotox (non- bact)	Repro/ develop
8050-09-7 Rosin	Test	Test	Adeq.	Test	Test	Test	Adeq.	Adeq.	Adeq.	Adeq.	Adeq Repro./ Test Develop.
65997-06-0, Rosin, hydrogenated	Test	Test	Test	С	С	С	Adeq.	Adeq.	Adeq.	Adeq.	С
68425-08-1, Rosin, distillation overheads	Test	Test	Test	С	С	С	С	С	С	С	O
68783-82-4 Rosin, low boiling fraction	No test	No test	No test	С	С	С	С	С	С	С	O
61790-50-9, Rosin, potassium salt	Test	Test	Test	С	С	С	С	С	С	С	С
61790-51-0, Rosin, sodium salt	Test	Test	Adeq.	С	С	С	С	С	С	С	С

Adeq. Indicates adequate existing data

**Test** Indicates proposed testing

NoTest See test plan; essentially identical to rosin, distillation overheads

C Indicates category read-down from existing or proposed test data on rosin.

\* No testing will be conducted for melting point, boiling point, vapor pressure, hydrolysis, photodegradation and transport and distribution between environmental compartments as explained in the test plan.

All the substances in this category are similar in chemical composition, being predominantly a mixture or resin acids or their salts. The selection of rosin as the substance to be tested is based on two factors. It has by far the greatest production volume. Production of rosin in the U.S. is almost four times higher than production of the other substances in this category combined. EPA guidance suggests that testing the substance produced at the greatest volume as the representative chemical of a category would be appropriate. Clearly, rosin fits this criterion. In addition, rosin is the raw material from which all the other category members are derived.

Another criterion listed by EPA for grouping chemicals into a category is the use of the "family approach" of examining related chemicals when they are acids or acid salts. Although the salts of rosin have quite different physical characteristics, they are included in this category because they are quickly converted into the free acids when they are neutralized by acid or by dilution, as they would be under typical toxicity testing conditions. In summary, this group of chemicals fits the requirements of the EPA's HPV Challenge program for a chemical category, and rosin is the most appropriate representative test material from this category.

### III. Review of Existing Data and Development of Test Plan

PCA has undertaken a comprehensive evaluation of all relevant data on the SIDS endpoints of concern for the chemicals in this category. Considerable data are available that satisfy most of the SIDS endpoints for this category. The availability of the data on the specific SIDS endpoints is summarized in Table 4 (identical to Table 1). Table 4 also shows data gaps that will be filled by additional testing, and areas where data from rosin and hydrogenated rosin will be generalized to other category members.

#### A. Evaluation of Existing Physicochemical Data and Proposed Testing

The basic physicochemical data required in the SIDS battery includes melting point, boiling point, vapor pressure, partition coefficient (K<sub>ow</sub>), and water solubility.

Class 2 substances are composed of a complex mixture of substances and are often difficult to characterize. Rosin and its derivatives are not only Class 2 substances, but also are derived from natural sources. Their composition is variable and cannot be represented by a single chemical structural diagram. Due to this "complex mixture" characteristic of rosin and related compounds, some physical property measurements, such as partition coefficient do not give singe definitive results because the methodology used to determine these properties will actually fractionate or partition the substance into various components. Since the methodology will alter the actual sample composition, the results are likely to be erroneous, difficult to interpret, or meaningless.

#### 1. Melting Point

Due to their complex nature, none of the members of this category have a well-defined melting point. These substances soften when heated and so have softening points rather than a true melting point. The softening point of these compounds can cover a wide range depending on the levels of resin acids, decarboxylated rosin and dimerized rosin in the sample, and hence these substances do not have specific softening points. The salts of rosin decompose on heating, and so melting point has no significance for these materials. Consequently, the melting point of these substances will not be measured.

#### 2. Boiling Point

All of the members of this category are produced by high temperature, high vacuum distillation and are non-volatile solids at ambient temperatures. A boiling point at ambient temperature has no significance because these materials will thermally decompose before they boil. Accordingly, measurement of this property is inappropriate for all the substances in this category.

#### 3. Vapor Pressure

Vapor pressures for the rosins (which are solids) at ambient temperatures are effectively zero, and their experimental measurement is inappropriate. When rosin salts are dissolved in water, their solutions will reflect the vapor pressure of the water rather than the salt, and therefore measurement of this property is inappropriate.

#### 4. Water Solubility

The water solubility of five of the compounds in this category will be determined using OECD (105).

#### 5. Partition Coefficient

The partition coefficient (i.e.,  $K_{ow}$ ) for five compounds in this category will be determined. Adequate data exist for rosin although it will be retested with the other compounds in this category. For rosin, existing data demonstrate that a range of  $K_{ow}$  values, rather than a single value, are generated when this endpoint is determined. This outcome reflects the complex nature of Class 2 mixtures.

Summary of Physicochemical Properties Testing: The water solubility and partition coefficients of five of the substances in this category will be determined. Adequate data on partition coefficient exist for rosin although it will be retested with the other compounds in this category. Tests for melting point, boiling point, and vapor pressure are inapplicable to these substances.

#### B. Evaluation of Existing Environmental Fate Data and Proposed Testing

The fate or behavior of a chemical in the environment is determined by the reaction rates for the most important transformation (degradation) processes. The basic environmental fate data covered by the HPV Program include biodegradation, stability in water (hydrolysis as a function of pH), photodegradation and transport and distribution between environmental compartments.

#### 1. Biodegradation

Biodegradability provides a measure for the potential of compounds to be degraded by microorganisms. Depending on the nature of the test material, several standard test methods are available to assess potential biodegradability.

Two of the chemicals in this category (rosin and the sodium salt of rosin) have existing data on the biodegradation endpoint. Biodegradation for hydrogenated rosin, rosin distillation overheads and the potassium salt will be determined using OECD method 302B for the salt and OECD method 301B for the non-salts.

#### 2. Hydrolysis

Hydrolysis as a function of pH is used to assess the stability of a substance in water. Hydrolysis is a reaction in which a water molecule (or hydroxide ion) substitutes for another atom or group of atoms present in an organic molecule. If there is no functional group suitable to be displaced, then the organic compound is considered to be resistant to hydrolysis. None of the substances in the rosin category contains an organic functional group that might be susceptible to this physical degradative mechanism. Therefore, hydrolysis need not be measured.

In addition, low water solubility often limits the ability to determine hydrolysis as a function of pH. All of the non-salt rosin compounds have very low solubility in water. Therefore, these materials are expected to be stable in water and it would be unnecessary to attempt to measure the products of hydrolysis. With respect to the rosin salts, in an aqueous medium they hydrolyze (ionize) immediately, but form stable species. Consequently, it would also be unnecessary to measure this endpoint for the rosin salts.

#### 3. Photodegradation

Due to their lack of any vapor pressure under ambient conditions, there is essentially no opportunity for any of these chemicals to enter the atmosphere. Thus, photodegradation is irrelevant. In addition, based on the constituents in these complex mixtures, there is no reason to suspect that they would be subject to breakdown by a photodegradative mechanism. Consequently, this endpoint will not be determined for any of the substances in this category.

#### 4. Transport and Distribution Between Environmental Compartments

The transport and distribution between environmental compartments is intended to determine the ability of a chemical to move or partition in the environment. The determination of this property requires the use of various models (e.g., level III model from the Canadian Environment Modeling Centre at Trent University). For Class 2 substances such as rosin and related compounds, the required inputs to the model are either not available or impossible to determine including molecular mass, reaction half-

life estimates for air, water, soil, sediment, aerosols, suspended sediment, and aquatic biota. In addition, while the partition coefficient is also required and can be determined, the multiple  $K_{ow}$  values typically derived for these substances (e.g., five  $K_{ow}$  values for rosin) are a consequence of sample fractionation and reflect various components in the mixture and are not representative of the mixture itself. Consequently, due to the inability to provide usable inputs to the required model, no determination of transportation and distribution between environmental compartments will be undertaken for rosin and related compounds.

Summary of Environmental Fate Testing: Biodegradation data will be generated for three of the compounds in this category for which data are not already available using OECD method 302B for the salt and OECD method 301B for the non-salts. Photodegradation, hydrolysis and transport and distribution between environmental compartments are not applicable to these chemicals.

#### C. Evaluation of Existing Ecotoxicity Data and Proposed Testing

The basic ecotoxicity data that are part of the HPV Program include acute toxicity to fish, daphnia and algae. While there are existing data on these endpoints for some of the substances in this grouping category, these data are conflicting and it is impossible to determine which, if any, of these findings is representative of true ecotoxicity. The inconsistencies in how water samples were prepared for testing these endpoints render these data inadequate. Consequently, acute toxicity to fish, daphnia and alga will be retested for rosin under conditions that maximize the solubility under the specific test exposure conditions, but reduce exposure to insoluble fractions, which may cause nonspecific toxicological effects. In addition, the effect of both filtering, to further minimize nonspecific physical effects, and of reducing the pH to the lower end of the acceptable range for test organism survival, will also be investigated for changes in toxicological effects. The results of preliminary tests will be used to select the most appropriate test conditions for the definitive test for each species.

Summary of Ecotoxicity Testing: The acute toxicity of rosin to fish, daphnia and algae will be tested under conditions that maximize solubility, but reduce exposure to insoluble fractions, which may cause nonspecific toxicological effects.

#### D. Evaluation of Existing Human Health Effects Data and Proposed Testing

#### 1. Acute Oral Toxicity

Acute oral toxicity studies investigate the effect(s) of a single exposure to a relatively high dose of a substance. This test is conducted by administering the test material to animals (typically rats or mice) in a single gavage dose. Harmonized EPA testing guidelines (August 1998) set the limit dose for acute oral toxicity studies at 2000 mg/kg body weight. If less than 50 percent mortality is observed at the limit dose, no further

testing is needed. A test substance that shows no effects at the limit dose is considered essentially nontoxic. If compound-related mortality is observed, then further testing may be necessary.

#### **Summary of Available Acute Oral Toxicity Data**

Both rosin and hydrogenated rosin are non-toxic following acute oral exposure. The acute oral  $LD_{50}$  values of various rosins (wood, gum and tall oil) are > 4000 mg/kg in rats, mice and guinea pigs. The acute oral  $LD_{50}$  value of hydrogenated rosin is > 32,000 mg/kg.

Summary of Acute Oral Toxicity Testing: Both rosin and hydrogenated rosin have been tested for acute oral toxicity and found to be non-toxic (i.e.,  $LD_{50} > 4000 \text{ mg/kg}$ ) well above the guideline of 2000 mg/kg. Consequently, additional testing for this endpoint is not necessary.

#### 2. Repeat Dose Toxicity

Subchronic repeat dose toxicity studies are designed to evaluate the effect of repeated exposure to a chemical over a significant period of the life span of an animal. Typically, the exposure regimen in a subchronic study involves daily exposure (at least 5 consecutive days per week) for a period of not less than 28 days or up to 90 days (i.e., 4 to 13 weeks). The HPV program calls for a repeat dose test of at least 28 days. The dose levels evaluated are lower than the relatively high doses used in acute toxicity (i.e., LD<sub>50</sub>) studies. In general, repeat dose studies are designed to assess systemic toxicity, but the study protocol can be modified to incorporate evaluation of potential adverse reproductive and/or developmental effects.

#### **Summary of Available Repeat Dose Toxicity Data**

There are existing data that demonstrate low toxicity for both rosin and hydrogenated rosin in repeat dose tests. Rosin was tested in a 90-day subchronic toxicity study in rats. The test material was administered to Sprague-Dawley rats in the diet at concentrations of 0, 0.01, 0.05, 0.20, 1.0 and 5.0% for 90 days. The approximate doses were 0, 10, 50, 200, 1000, or 5000 mg/kg/day. Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, hematology, clinical chemistry, urinalysis, gross and microscopic pathology, and organ weights.

All animals in the 5% dose group died within a week due to palatability issues resulting in complete cessation of food consumption. Some animals in the 1% dose group also failed to gain weight compared to controls due to decreased food consumption that resulted in some decreased organ weight to body weight ratios. No changes in hematology, clinical chemistry or urinalysis parameters were measured at any dose level. At gross pathology, no treatment-related effects were noted. No consistent organ weight changes and no histopathological effects were reported. Based on these data, the No Observed Effect Level (NOEL) was 0.20% (approximately 200 mg/kg/day).

Other 90-day subchronic studies confirm the low toxicity of rosin. In these studies, the only effect noted was either death due to palatability resulting in non-consumption of food or depression of body weight gain at the highest doses tested. In a dietary study with hydrogenated rosin, weanling Sprague-Dawley rats were exposed at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1000, or 5000 mg/kg/day. Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food utilization, hematology parameters, urinalysis, organ weights and gross and microscopic pathology.

All the animals in the high-dose group died prior to study termination due to treatment-related starvation through food refusal. In the 1% group, body weight was significantly decreased in both males and females, and food consumption was decreased. Food utilization was not affected at a dietary concentration of 1% indicating that the reduced food consumption was related to palatability. No treatment-related effects on hematology, urinalysis, or gross or microscopic pathology. Based on these data, the NOEL was 0.2% (approximately 200 mg/kg/day).

Summary of Repeat Dose Toxicity Testing: Rosin and hydrogenated rosin have been tested for repeat dose toxicity in 90-day studies. In these studies, the NOELs for both rosin and hydrogenated rosin were approximately 200 mg/kg/day, indicating that these compounds have low toxicity. Additional studies on these compounds support this result. Consequently, no additional testing for this endpoint will be conducted.

#### 3. Genotoxicity - In vitro

Genetic testing is conducted to determine the effects of substances on genetic material (i.e., DNA and chromosomes). The gene, which is composed of DNA, is the simplest functional genetic unit. Mutations of genes can occur spontaneously or as a consequence of exposure to chemicals or radiation. Genetic mutations are commonly measured in bacterial and mammalian cells, and the HPV program calls for completing both types of tests.

#### **Summary of Available Genotoxicity Data**

Rosin and hydrogenated rosin have been tested for potential carcinogenicity in several two-year bioassays conducted in rats. None of these studies demonstrated any evidence of carcinogenicity. The primary effect was depressed weight gain at the highest dose, confirming that a maximally tolerated dose was achieved.

Since the purpose of *in vitro* bacterial and mammalian mutagenicity tests is to determine if a chemical might have the potential to be a direct-acting DNA reactive carcinogen, the negative carcinogenicity studies eliminate the need to test for potential genotoxicity.

Summary of Genotoxicity Testing: Neither rosin nor hydrogenated rosin were carcinogenic when tested in two-year cancer bioassays. Consequently, no genotoxicity testing is necessary.

#### 4. Reproductive and Developmental Toxicity

Reproductive toxicity includes any adverse effect on fertility and reproduction, including effects on gonadal function, mating behavior, conception, and parturition. Developmental toxicity is any adverse effect induced during the period of fetal development, including structural abnormalities, altered growth and post-partum development of the offspring.

The "toxicity to reproduction" aspect of the HPV Challenge Program can be met by conducting a reproductive/developmental toxicity screening test or adding a reproductive/developmental toxicity screening test to the repeat dose study (OECD 421 or OECD 422, respectively). The one-generation reproduction toxicity study (OECD 415) is a more comprehensive protocol for the study of the effect of a test material on reproduction and development that also meets the SIDS and the HPV Program requirements.

#### **Summary of Reproductive/Developmental Toxicity Data**

As noted in the SIDS guidelines for the reproduction toxicity endpoint, "when a 90-day repeated dose study is available and demonstrates no effects on the reproductive organs, in particular the testes, then a developmental study can be considered as an adequate test to complete information on reproduction/developmental effect." Rosin and hydrogenated rosin have been tested in 90-day repeat dose studies as well as in two-year bioassays. Both types of studies included histopathology of reproductive organs (i.e., testes, ovaries, uterus) and showed no evidence of reproductive organ toxicity at any dose level. Therefore, these studies satisfy the SIDS reproductive toxicity endpoint. However, since neither of these studies evaluated the developmental toxicity endpoint, this will be determined on rosin using OECD protocol 421.

Summary of Reproductive/Developmental Testing: Neither rosin nor hydrogenated rosin demonstrated any effects on reproductive toxicity in numerous repeat dose studies. However, since none of these studies evaluated potential developmental toxicity, rosin will be tested for this endpoint with OECD protocol 421.

## References

EPA. 2000. Data Collection and Development on High Production Volume (HPV) Chemicals. Fed. Reg. Dec. 26, Vol. 65(248): pp. 81686-81698.

Zinkel, D.F. and Russell, J., Eds. 1989. Naval Stores. Production, Chemistry, Utilization. Pulp Chemicals Association, New York.

September 2001

## **Robust Summaries of Existing Data**

PHYSICO-CHEMICAL PROPERTY – OCTANOL/WATER PARTITION COEFFICIENT			
Test Substance			
Chemical Name	Tall oil rosin		
CAS #	8050-09-7		
Remarks			
Method	test plan for Rosins and Rosin Salts.		
Method/Guideline	Testing was conducted according to OECD Test Method		
followed	117, "Partition Coefficient (n-Octanol/Water) High		
	Performance Liquid Chromatograph (HPLC) Method"		
Test Type	Partition coefficient		
GLP (Y/N)	Υ		
Year (Study Performed)	1993		
Test conditions	Tall oil rosin was dissolved in methanol and the solution was analyzed by HPLC with UV detection using a mobile phase of methanol:buffer (3:1) at pH 2 and pH 7.5. As a reference substance, a mixture of seven materials was used.		
Results	At pH 2, the log $P_{ow}$ [K <sub>ow</sub> ] values of five components in tall oil rosin were 4.5, 6.1, 6.9, 7.1, and 7.2. At pH 7.5, the log $P_{ow}$ value of one component in tall oil rosin was 3.6.		
<u>Data Quality</u>	Reliable without restrictions – Klimisch Code 1a Note: the various K <sub>ow</sub> values reflect the components in the mixture and not the mixture <i>per se</i> .		
<u>References</u>	Dybdahl, H.P. 1993. Determination of log P <sub>ow</sub> for single components in tall oil rosin. GLP Study No. 408335/471. Water Quality Institute, Horshplm, Denmark.		

Test Substance	
Chemical Name	Tall oil rosin
CAS #	8050-09-7
Remarks	This substance is referred to as tall oil rosin or rosin in the test plan for Rosins and Rosin Salts.
<u>Method</u>	
Method/Guideline followed	Testing was conducted according to OECD Test Method 301 D, "Ready Biodegradability: Closed Bottle Test"
Test Type (aerobic/anaerobic)	Aerobic
GLP (Y/N)	Υ
Year (Study Performed)	1993
Contact time	28 days
Inoculum	Secondary effluent from Rungsted Treatment plant
Test conditions	Inoculum: Secondary effluent was collected from Rungsted Treatment plant.
	Concentration of test chemical: A stock solution of the test material (2 g/L) was prepared in demineralized water by ultra sonication for 5 minutes and magnetic stirring for 24 hours at 20°C. The solution was filtered and after determination of the chemical oxygen demand was used within 1 day.
	Test Setup: Test medium was prepared by adding 1 mL each of four solutions (potassium phosphate, magnesium sulfate, calcium chloride, ferric chloride) to 1 liter of demineralized water, which was aerated to an initial oxygen concentration of approximately 9 mg O <sub>2</sub> /L and inoculated with 1 drop of secondary effluent per liter. The test article was added at 0.21 g/L to a part of the inoculated test medium, equivalent to a chemical oxygen demand of 4.52 mg O <sub>2</sub> /L. Sodium benzoate, the reference compound, was added at 2 mg/L to another part of the inoculated medium (to assess the activity of the inoculum), equivalent to a theoretical oxygen demand of 3.34 mg O <sub>2</sub> /L. Both the test and reference articles (0.21 g/L and 2 mg/L) were added to a third part of the inoculated medium (to assess possible inhibitory effects of the test article), at a theoretical oxygen demand of 7.86 mg O <sub>2</sub> /L. Blank controls were prepared using the inoculated medium without test or reference materials. After the samples were prepared, the medium was transferred to calibrated respirometric bottles (BOD bottles), and placed in the dark at 20°C. The study was performed in triplicate.

	Sampling frequency: Samples were collected for BOD analysis on days 0, 7, 14, 21, and 28.
	Controls: Yes.
	Method of calculating oxygen demand: Oxygen demand was calculated as the difference between the measured oxygen concentrations at time t and the start of the test. Biological oxygen demand for the added carbon sources was calculated by subtracting the oxygen demand for the blank controls from the oxygen demand in the bottles containing test and reference compounds.
<u>Results</u>	
Degradation % after time	23% after 7 days and 32% after 28 days (test article); 59% after 7 days and 88% after 28 days (sodium benzoate)
Conclusions	The biological oxygen demand for tall oil rosin was 23 and 32% of the theoretical oxygen demand after 7 and 28 days, respectively. These data indicate that the material is dominated by recalcitrant compounds. Tall oil rosin did not inhibit the respiratory activity of the inoculum. The inoculum had satisfactory activity as demonstrated by 60% degradation within the 7 days using the reference compound.
Data Quality	Reliable without restrictions- Klimisch Code 1a
<u>Reference</u>	Madsen, T. 1993. Biodegradation of tall oil rosin. GLP Study No. 308067/471. Water Quality Institute, Horshplm, Denmark.

Test Substance	
Chemical Name	Rosin, sodium salt
CAS #	61790-51-0
Remarks	This substance is referred to as the sodium salt of rosin in the test plan for Rosins and Rosin Salts.
<u>Method</u>	•
Method/Guideline followed	Testing was conducted using the Shake Flask method similar to OECD Test Method 307.
Test Type	Aerobic
(aerobic/anaerobic)	
GLP (Y/N)	N
Year (Study Performed)	1965
Contact time	32 days
Inoculum	Activated sludge from the Bergen County Sewage Authority treatment plant in Little Ferry, N.J.
Test conditions	Inoculum: Activated sludge from the Bergen County Sewage Authority treatment plant in Little Ferry, N.J.
	Concentrations of test and reference chemicals: The test and reference chemicals were used at a concentration of 50 ppm.
	Test Setup: Test medium consisted of magnesium nitrate, calcium nitrate, ferric nitrate, calcium nitrate, cobaltous chloride, diammonium hydrogen phosphate, dipotassium hydrogen phosphate, and monopotassium hydrogen phosphate all dissolved in distilled water. A blank unit (containing all nutrients except the test materials) was treated in the same manner. Microbial cultures were added at a concentration of 10 mg/l on a dry-weight bases to begin the tests. All solutions were placed in Erlenmeyer flasks that were mounted on a shaker for aeration. The study was performed in triplicate.
	Sampling frequency: Samples were collected for determination of chemical oxygen demand (COD) on an almost daily basis.
	Controls: Yes. Linear alkylbenzene sulfonate (LAS)
	Method of calculating chemical oxygen demand: COD was calculated as the difference between the measured oxygen concentrations at various sampling times and the start of the test. COD for the samples was calculated by subtracting the COD for the blank controls from the COD in the flasks containing test and reference compounds.

Results	
Degradation % after time	70-80% after 21 days (test article) and 97% after 21 days
	(reference compound)
<u>Conclusions</u>	These data indicate that the sodium salt of rosin is readily
	biodegradable.
Data Quality	Reliable with restrictions– Klimisch Code 2e
<u>Reference</u>	Eldib, I.A. 1965. Biodegradability evaluation of (trade name deleted) [rosin, sodium salt]. Eldib Engineering and Research, Newark, N.J.

ACUTE TOXICITY – ORAL				
Test substance				
Chemical Name	Rosin			
CAS #	8050-09-7			
Remarks	This substance is referred to as rosin in the test plan for			
	Rosins and Rosin Salts.			
<u>Method</u>				
Method/Guideline followed	Test procedure was similar to OECD Test Method 401, "Acute Oral Toxicity"			
GLP (Y/N)	N			
Year (Study Performed)	1961			
Species	Rats, mice, guinea pigs			
Strain	Not specified			
Route of administration	Oral			
Dose levels	Dose levels not specified.			
Sex and number/group	10 male rats, mice or guinea pigs			
Frequency of treatment	Single oral gavage			
Duration of test	14 day observation post-treatment			
Control group (Y/N)	N			
<u>Results</u>				
Acute Oral LD <sub>50</sub>	Rats: 7,600, 8,400, and 7,600 mg/kg for gum, wood and tall oil rosin, respectively; mice: 4,600, 4,100 and 4,600 mg/kg for gum, wood and tall oil rosin, respectively; guinea pigs: 4,100, 4,100 and 4,600 mg/kg for gum, wood and tall oil rosin, respectively.			
<u>Detailed Summary</u>	Male rats, mice or guinea pigs (n = 10) received graded oral gavage doses in corn oil of gum, wood or tall oil rosin (CAS #8050-09-7) and were observed for 14 days. Parameters evaluated included clinical signs, mortality, and gross necropsy. The oral $LD_{50}$ values were calculated according to the method of Litchfield and Wilcoxon.			
Data Quality	Reliable with restriction – Klimisch Code 2c			
<u>Reference</u>	Kay, J.H. 1961. Acute toxicity of rosins. Industrial Bio-Test Laboratories, Northbrook, IL.			

ACUTE TOXICITY – ORAL			
Test substance			
Chemical Name	Hydrogenated rosin		
CAS#	65997-06-0		
Remarks	This substance is referred to as hydrogenated rosin in the test plan for Rosins and Rosin Salts.		
<u>Method</u>			
Method/Guideline followed	Test procedure was similar to OECD Test Method 401, "Acute Oral Toxicity"		
GLP (Y/N)	N		
Year (Study Performed)	1982		
Species	Rat		
Strain	Wistar		
Route of administration	Oral		
Dose levels	30 mL/kg (approximately equivalent to 32,000 mg/kg, based on a density of 1.05 g/mL)		
Sex and number/group	10 male and 10 female rats		
Frequency of treatment	Single oral gavage		
Duration of test	14 day observation post-treatment		
Control group (Y/N)	N		
<u>Result</u>			
Acute Oral LD <sub>50</sub>	>30 mL/kg (or 32,000 mg/kg)		
<u>Detailed Summary</u>	Wistar rats (n = 10/sex) received a single oral dose of 30 mL/kg hydrogenated rosin (CAS #65997-06-0) and were observed for 14 days. Parameters evaluated included clinical signs, mortality, and gross necropsy. One day after dosing, the rats were sluggish and had slight diarrhea. All of the animals recovered during the observation period and no deaths occurred. Gross necropsy revealed no treatment-related effects; the oral LD $_{50}$ was reported as greater than 30 mL/kg (approximately equivalent to 32 g/kg, based on a density of 1.05 g/mL).		
<u>Data Quality</u>	Valid with restriction – Klimisch Code 1b		
<u>Reference</u>	Spanjers, M.Th. 1981. Determination of the acute oral toxicity of [hydrogenated rosin trade name deleted] in rats. CIVO-TNO.		

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS #	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
T to marke	Rosins and Rosin Salts.
Method	
Method/Guideline followed	Test procedure is similar to OECD Test Method 407, "Repeat Dose 28-Day Oral Toxicity Study in Rodents."
Year	1960
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	90 days
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.01, 0.05, 0.2, 1, or 5%
Control group (Y/N)	Υ
Results	
NOEL:	0.2%, approximately 200 mg/kg/day
<u>Detailed Summary</u>	Weanling Sprague-Dawley rats (n = 10/sex/group) were treated with gum rosin (CAS # 8050-09-7) in the diet at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, hematocrit, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes/ovaries), and microscopic pathology (brain, liver, spleen, stomach, small intestine, colon, pancreas, kidneys, urinary bladder, adrenals, gonads, thyroid, parathyroid, lymph nodes, heart, lungs, bone marrow, muscle, prostate, uterus).
	All of the rats dosed with 5% gum rosin died between days 3 and 7. These animals exhibited significant weight loss and a marked decrease in food consumption. Starvation through refusal to eat was considered the primary cause of death. No other mortalities occurred and no adverse clinical signs were noted. A decrease in mean body weight was reported in rats treated with 1%; body weight gain was also slightly decreased in this group. Food consumption

	and food utilization (grams gained/grams food consumed)
	were decreased at 1%. For food utilization, the decrease
	occurred during the first two weeks, but was comparable to
	control levels thereafter indicating that palatability was the
	principal cause of the depression. The body weight and
	food effects were primarily noted in the first few weeks of
	the study. No treatment-related effects on hematology or
	urinalysis parameters were reported. At necropsy, no
	changes were noted that were related to treatment.
	Absolute organ weights were not affected, but some of the
	relative weights in the 1% group were altered. These
	changes were related to the decreased body weight in this
	group and were not considered to be a direct treatment
	effect. No histopathological changes were observed in any
	organ. Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus)
	showed no evidence of toxicity at any dose level. Based on
	these data, it appears that the NOEL was 0.2%
D. ( ) O . ( ) ( )	(approximately 200 mg/kg/day).
Data Quality	Valid without restriction – Klimisch Code 1b
<u>References</u>	Calandra, J.C. 1960. Ninety-day subacute oral toxicity of
	gum rosin. Industrial Bio-Test Laboratories, Inc.,
	Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for
	the Toxicological Assessment of Pesticide Residues in
	Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS #	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
	Rosins and Rosin Salts.
<u>Method</u>	
Method/Guideline followed	Test procedure is similar to OECD Test Method 407,
	"Repeat Dose 28-Day Oral Toxicity Study in Rodents."
Year	1960
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	90 days
Frequency of Treatment	Daily
Post-exposure observation	None
period	0.004.005.004.550/
Dose Levels	0, 0.01, 0.05, 0.2, 1, or 5%
Control group (Y/N)	Υ
Results	0.00/
NOEL: Detailed Summary	0.2%, approximately 200 mg/kg/day Weanling Sprague-Dawley rats (n = 10/sex/group) were
	treated with rosin (CAS # 8050-09-7) in the diet at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, hematocrit, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes/ovaries), and microscopic pathology (brain, liver, spleen, stomach, small intestine, colon, pancreas, kidneys, urinary bladder, adrenals, gonads, thyroid, parathyroid, lymph nodes, heart, lungs, bone marrow, muscle, prostate, uterus).
	All animals treated with 5% rosin died between days 4 and 12. These animals exhibited significant weight loss and a marked decrease in food consumption. Death was related to starvation associated with food refusal. One other death occurred on day 77 in the low-dose group, but this was not treatment-related. No adverse clinical signs were noted in any group. Decreases in mean body weight and body weight gain were reported in rats treated with 1%. In

	addition, food consumption and food utilization (grams gained/grams food consumed) were decreased at this dose level. For food utilization, the decrease primarily occurred during the first week of dosing, but was comparable to control levels thereafter indicating that palatability was the principal cause of the depression. A slight decrease in body weight, body weight gain and food consumption were reported at 0.2%; food utilization was unaffected in this group. No treatment-related effects on hematology or urinalysis parameters were reported. Statistical analyses
	revealed increases in absolute liver weights in the 1% animals as well as increases in select organ to body weight ratios (primarily liver, kidney, spleen in males, and liver in females). These changes were not considered to be toxicologically significant because the animals in this group exhibited decreased body weight and no histopathological changes were observed in any organ. Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus) showed no evidence of toxicity at any dose level. Based on these data, it appears that the NOAEL was 0.2% (approximately 200 mg/kg/day).
Data Quality	Valid without restriction – Klimisch Code 1b
References	Calandra, J.C. 1960. Ninety-day subacute oral toxicity of [trade name deleted] rosin. Industrial Bio-Test Laboratories, Inc., Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for the Toxicological Assessment of Pesticide Residues in Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS #	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for Rosins and Rosin Salts.
<u>Method</u>	
Method/Guideline followed	Test procedure is similar to OECD Test Method 407, "Repeat Dose 28-Day Oral Toxicity Study in Rodents."
Year	1960
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	90 days
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.01, 0.05, 0.2, 1, or 5%
Control group (Y/N)	Υ
<u>Results</u>	
NOAEL:	0.2%, approximately 200 mg/kg/day
Detailed Summary	Weanling Sprague-Dawley rats (n = 10/sex/group) were treated with rosin (CAS # 8050-09-7) in the diet at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, hematocrit, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes/ovaries), and microscopic pathology (brain, liver, spleen, stomach, small intestine, colon, pancreas, kidneys, urinary bladder, adrenals, gonads, thyroid, parathyroid, lymph nodes, heart, lungs, bone marrow, muscle, prostate, uterus).
	Eighteen of the animals treated with 5% resin died between days 4 and 18. An additional two animals died on days 46 and 77. These animals exhibited significant weight loss and decreases in food consumption and food utilization. Death was related to starvation associated with food refusal. One death occurred on day 70 at 0.01% and another death occurred on day 54 at 0.2%; these were isolated findings and were not considered to be treatment-

	,
	related. No adverse clinical signs were noted in any group.
	Decreases in mean body weight and body weight gain were
	reported in rats treated with 0.2 and 1%; the effect was
	slight at 0.2%. Food consumption and food utilization were
	decreased at 1%. No treatment-related effects on
	hematology or urinalysis parameters were reported. At
	necropsy, no adverse effects were reported. Some organ
	weight alterations were noted in the 1% dose group, but
	due to the significant depression in body weights in this
	group, the effects are not considered to be toxicologically
	significant. No histopathological changes were observed in
	any organ. Reproductive organs (i.e., testes, ovaries,
	uterus) showed no evidence of toxicity at any dose level.
	Based on these data, it appears that the NOAEL was 0.2%
	(approximately 200 mg/kg/day).
Data Quality	Valid without restriction – Klimisch Code 1b
<u>References</u>	Calandra, J.C. 1960. Ninety-day subacute oral toxicity of
	resin [trade name deleted]. Industrial Bio-Test
	Laboratories, Inc., Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for
	the Toxicological Assessment of Pesticide Residues in
	Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS#	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
	Rosins and Rosin Salts.
Method	
Method/Guideline followed	Test procedure is similar to OECD Test Method 407, "Repeat Dose 28-Day Oral Toxicity Study in Rodents."
Year	1960
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	90 days
Frequency of Treatment	Daily
Post-exposure observation	None
period	Notic
Dose Levels	0, 0.01, 0.05, 0.2, 1, or 5%
Control group (Y/N)	Υ
Results	
NOEL:	0.2%, approximately 200 mg/kg/day
Detailed Summary	Weanling Sprague-Dawley rats (n = 10/sex/group) were treated with wood rosin (CAS # 8050-09-7) in the diet at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO (1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, erythrocyte count, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes/ovaries), and microscopic pathology (brain, liver, spleen, stomach, small intestine, colon, pancreas, kidneys, urinary bladder, adrenals, gonads, thyroid, parathyroid, lymph nodes, heart, lungs, bone marrow, muscle, prostate, uterus).  Nine rats treated with 5% wood rosin died on days 7 and 8; the deaths were related to starvation associated with food refusal. No other mortalities occurred during the study, and no adverse clinical signs were noted in any group. The animals treated with 5% exhibited significant weight loss, especially during the first six weeks of the study. A marked decrease in food consumption was also noted in the high-dose animals, but this corresponded more to food

	"disappearance" (i.e., scattering) than to a true decrease in
	food consumption; palatability was considered to be the
	primary factor involved. Decreases in mean body weight
	and body weight gain were reported in rats treated with 1%,
	but the decreases were not statistically significant. Food
	consumption was not affected in the 1% group, but food
	utilization was slightly decreased. No treatment-related
	effects on hematology or urinalysis parameters were
	reported. At necropsy, the kidneys of the high-dose
	animals (5% group) were described as: stippled and yellow
	in color; the cortex was thin; the cortico-medullary junction
	was indistinct; and there were discolored patches
	intermingled with cyst-like areas. These effects were not
	observed in any other dose group. Significant increases in
	absolute and relative liver weights were reported at 1 and
	5%. Other organ weight changes noted at 5% were
	associated with the depression in body weight in this group.
	Histopathological examination revealed marked dilation and
	tortuosity of the renal distal convoluted and collecting
	segment tubules of the high-dose rats. In addition, a few
	glomeruli revealed active inflammatory and degenerative
	changes without proliferation or organization. Reproductive
	organs ( <i>i.e.</i> , testes, ovaries, uterus) showed no evidence of
	toxicity at any dose level. Based on these data, it appears
	that the NOEL was 0.2% (approximately 200 mg/kg/day).
Data Quality	Valid without restriction – Klimisch Code 1b
<u>References</u>	Calandra, J.C. 1960. Ninety-day subacute oral toxicity of
	b-wood resin. Industrial Bio-Test Laboratories, Inc.,
	Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for
	the Toxicological Assessment of Pesticide Residues in
	Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS #	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
T to marke	Rosins and Rosin Salts.
Method	
Method/Guideline followed	Test procedure is similar to OECD Test Method 407, "Repeat Dose 28-Day Oral Toxicity Study in Rodents."
Year	1960
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	90 days
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.01, 0.05, 0.2, 1, or 5%
Control group (Y/N)	Υ
Results	
NOAEL:	0.2%, approximately 200 mg/kg/day
<u>Detailed Summary</u>	Weanling Sprague-Dawley rats (n = 10/sex/group) were treated with wood rosin (CAS # 8050-09-7) in the diet at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, hematocrit, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes/ovaries), and microscopic pathology (brain, liver, spleen, stomach, small intestine, colon, pancreas, kidneys, urinary bladder, adrenals, gonads, thyroid, parathyroid, lymph nodes, heart, lungs, bone marrow, muscle, prostate, uterus).
	All animals treated with 5% wood rosin died between days 3 and 8. These animals exhibited significant weight loss and a marked decrease in food consumption. Starvation through food refusal was identified as the primary cause of death. No adverse clinical signs were noted in any group. Slight decreases in mean body weight and body weight gain were reported in the rats treated with 1%. Food consumption was also slightly decreased at 1%, but food

	utilization was unaffected. In the 0.05% dose group, food
	consumption was decreased, but no explanation for this
	·
	"apparent discrepancy" was provided; this effect is unlikely
	to be treatment-related. No treatment-related effects on
	hematology or urinalysis parameters were reported. At
	necropsy, no gross changes were observed. Statistical
	analyses revealed increases in absolute liver weights as
	well as increases in liver to body and brain weight ratios in
	the 1% dose group. These changes were not considered
	to be toxicologically significant because the animals in this
	group exhibited decreased body weight and no
	histopathological changes were observed in any organ.
	Reproductive organs (i.e., testes, ovaries, uterus) showed
	no evidence of toxicity at any dose level. Based on these
	data, it appears that the NOAEL was 0.2% (approximately
	200 mg/kg/day).
<u>Data Quality</u>	Valid without restriction – Klimisch Code 1b
<u>References</u>	Calandra, J.C. 1960. Ninety-day subacute oral toxicity of
	n-wood rosin. Industrial Bio-Test Laboratories, Inc.,
	Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for
	the Toxicological Assessment of Pesticide Residues in
	Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS#	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
T to marke	Rosins and Rosin Salts.
Method	
Method/Guideline followed	Test procedure is similar to EPA Test Method OPPTS 870.4200, "Carcinogenicity."
Year	1962
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	Two years
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.05, 1%
Control group (Y/N)	Υ
<u>Results</u>	
NOEL:	0.05%, approximately 50 mg/kg/day
<u>Detailed Summary</u>	Weanling Sprague-Dawley rats (n = 30 /sex/dose) were exposed to gum rosin (CAS # 8050-09-7) at dietary concentrations of 0, 0.05, or 1% for two years. The approximate doses were 0, 50, or 1000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, erythrocyte count, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (liver, kidneys, spleen, gonads, heart, brain, thyroid gland, adrenal gland), tumor incidence (organs examined not specified), and microscopic pathology (heart, lung, trachea, liver, pancreas, stomach, small intestine, colon, spleen, lymph node, kidney, urinary bladder, testis, ovary, prostate, uterus, pituitary, adrenal gland, salivary gland, thyroid gland, parathyroid gland, skeletal muscle, bone marrow, brain).
	No treatment-related increase in mortality was reported. The only clinical signs were generalized inactivity and weakness in the animals dying on study. Mean body weight and body weight gain were statistically significantly decreased at 1%. Food consumption was also decreased in the high-dose group, but food utilization was only slightly

	decreased. These effects were attributed to the palatability of the test diet. No treatment-related effects were reported on hematology, urinalysis, organ weights, and gross and microscopic pathology. Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus) showed no evidence of toxicity at any dose level. The tumor incidence and tumor types were similar in the test and control groups. Based on these data, it appears that the NOEL was 0.05% (approximately 50 mg/kg/day).
<u>Data Quality</u> <u>References</u>	Valid without restriction – Klimisch Code 1b  Kay, J.H. 1962. Two-year chronic oral toxicity of b-wood resin – albino rats. Industrial Bio-Test Laboratories, Inc., Northbrook, Illinois.  World Health Organization (WHO). 1990. Principles for the Toxicological Assessment of Pesticide Residues in Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS #	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
	Rosins and Rosin Salts.
<u>Method</u>	
Method/Guideline followed	Test procedure is similar to EPA Test Method OPPTS 870.4200, "Carcinogenicity."
Year	1962
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	Two years
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.05, 0.2, 1%
Control group (Y/N)	Υ
Results	
NOEL:	0.2%, approximately 200 mg/kg/day
Detailed Summary	Weanling Sprague-Dawley rats (n = 30 /sex/dose) were exposed to wood rosin (CAS # 8050-09-7) at dietary concentrations of 0, 0.05, 0.2, or 1% for two years. The approximate doses were 0, 50, 200, or 1000 mg/kg/day, based on standard conversion factors provided by WHO (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, erythrocyte count, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (liver, kidneys, spleen, gonads, heart, brain, thyroid gland, adrenal gland), tumor incidence (organs examined not specified), and microscopic pathology (heart, lung, trachea, liver, pancreas, stomach, small intestine, colon, spleen, lymph node, kidney, urinary bladder, testis, ovary, prostate, uterus, pituitary, adrenal gland, salivary gland, thyroid gland, parathyroid gland, skeletal muscle, bone marrow, brain).
	No treatment-related increase in mortality was reported. The only clinical signs were generalized inactivity and weakness in the animals dying on study. In the high-dose group, mean body weight and body weight gain were statistically significantly decreased in both sexes at the

	interim (12-month) sacrifice and in the females at the
	terminal sacrifice. These parameters were slightly
	decreased in the males at 24 months. Food consumption
	was decreased in the high-dose group, but food utilization
	was generally comparable to control levels. The effects on
	body weight and food consumption were attributed to the
	palatability of the test diet. No treatment-related effects
	were reported on hematology, urinalysis, and gross and
	microscopic pathology. Relative liver weight was
	significantly increased in the females treated with 1%.
	Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus) showed
	no evidence of toxicity at any dose level. The tumor
	incidence and tumor types were similar in the test and
	control groups. Based on these data, it appears that the
	NOEL was 0.2% (approximately 200 mg/kg/day).
Data Ovality	
Data Quality	Valid without restriction – Klimisch Code 1b
<u>References</u>	Kay, J.H. 1962. Two-year chronic oral toxicity of n-wood
	rosin – albino rats. Industrial Bio-Test Laboratories, Inc.,
	Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for
	the Toxicological Assessment of Pesticide Residues in
	Food.
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REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Rosin
CAS#	8050-09-7
Remarks	This substance is referred to as rosin in the test plan for
T to marke	Rosins and Rosin Salts.
Method	
Method/Guideline followed	Test procedure is similar to EPA Test Method OPPTS 870.4200, "Carcinogenicity."
Year	1962
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	Two years
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.05, 1%
Control group (Y/N)	Υ
<u>Results</u>	
NOEL:	0.05%, approximately 50 mg/kg/day
<u>Detailed Summary</u>	Weanling Sprague-Dawley rats (n = 30 /sex/dose) were exposed to gum rosin (CAS # 8050-09-7) at dietary concentrations of 0, 0.05, or 1% for two years. The approximate doses were 0, 50, or 1000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, erythrocyte count, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (liver, kidneys, spleen, gonads, heart, brain, thyroid gland, adrenal gland), tumor incidence (organs examined not specified), and microscopic pathology (heart, lung, trachea, liver, pancreas, stomach, small intestine, colon, spleen, lymph node, kidney, urinary bladder, testis, ovary, prostate, uterus, pituitary, adrenal gland, salivary gland, thyroid gland, parathyroid gland, skeletal muscle, bone marrow, brain).
	No treatment-related increase in mortality was reported. The only clinical signs were generalized inactivity and weakness in the animals dying on study. Mean body weight and body weight gain were statistically significantly decreased at 1%. Food consumption was also decreased in the high-dose group, but food utilization was unaffected.

	The effects on body weight and food consumption were attributed to the palatability of the test diet. No treatment-related effects were reported on hematology, urinalysis, and gross and microscopic pathology. Relative liver weight was significantly increased at 1%. Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus) showed no evidence of toxicity at any dose level. The tumor incidence and tumor types were similar in the test and control groups. Based on these data, it appears that the NOEL was 0.05% (approximately 50 mg/kg/day).
<u>Data Quality</u>	Valid without restriction – Klimisch Code 1b
References	Kay, J.H. 1962. Two-year chronic oral toxicity of gum rosin – albino rats. Industrial Bio-Test Laboratories, Inc., Northbrook, Illinois.  World Health Organization (WHO). 1990. Principles for the Toxicological Assessment of Pesticide Residues in Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Hydrogenated rosin
CAS #	65997-06-0
Remarks	This substance is referred to as hydrogenated rosin in the test plan for Rosins and Rosin Salts.
<u>Method</u>	
Method/Guideline followed	Test procedure is similar to OECD Test Method 407, "Repeat Dose 28-Day Oral Toxicity Study in Rodents."
Year	1960
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	90 days
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.01, 0.05, 0.2, 1, or 5%
Control group (Y/N)	Υ
<u>Results</u>	
NOAEL:	0.2%, approximately 200 mg/kg/day
Detailed Summary	Weanling Sprague-Dawley rats (n = 10/sex/group) were treated with hydrogenated rosin (CAS # 65997-06-0) in the diet at concentrations of 0, 0.01, 0.05, 0.2, 1, or 5% for 90 days. The approximate doses were 0, 10, 50, 200, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food utilization, hematology parameters (hemoglobin concentration, hematocrit, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes/ovaries), and microscopic pathology (brain, liver, spleen, stomach, small intestine, colon, pancreas, kidneys, urinary bladder, adrenals, gonads, thyroid, parathyroid, lymph nodes, heart, lungs, bone marrow, muscle, prostate, uterus).
	All the animals in the high-dose group died prior to study termination. These deaths occurred between study day 3 and 11 and were attributed to starvation through food refusal ( <i>i.e.</i> , treatment-related). Rats in this group also experienced weight loss and a marked decrease in food consumption. In the 1% group, body weight was significantly decreased in both males and females, and food consumption was decreased. With the exception of the first week of dosing, food utilization (grams gained/gram

	food consumed) was not affected at a dietary concentration of 1% indicating that the reduced food consumption was related to its palatability. No treatment-related effects on hematology, urinalysis, or gross or microscopic pathology. Some statistically significant organ weight effects were reported in the 1% dose group, but these were considered to arise secondary to decreased body weight. Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus) showed /no evidence of toxicity at any dose level. Based on these data, it appears that the NOEL was 0.2% (approximately 200 mg/kg/day).
Data Quality References	Valid without restriction – Klimisch Code 1b Calandra, J.C. 1960. Ninety-day subacute oral toxicity of
110.0.01000	[trade name deleted] hydrogenated rosin. Industrial Bio- Test Laboratories, Inc., Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for the Toxicological Assessment of Pesticide Residues in Food.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Hydrogenated rosin
CAS #	65997-06-0
Remarks	This substance is referred to as hydrogenated rosin in the test plan for Rosins and Rosin Salts.
<u>Method</u>	
Method/Guideline followed	Test procedure is similar to EPA Test Method OPPTS 870.4200, "Carcinogenicity."
Year	1962
GLP (Y/N)	N (pre-GLP)
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	Two years
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0, 0.05, 0.2, 1%
Control group (Y/N)	Υ
<u>Results</u>	
NOEL: <b>Detailed Summary</b>	0.2%, approximately 200 mg/kg/day Weanling Sprague-Dawley rats (n = 30 /sex/dose) were
	exposed to hydrogenated rosin (CAS # 65997-06-0) at dietary concentrations of 0, 0.05, 0.2, or 1% for two years. The approximate doses were 0, 50, 200, or 1000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, mortality, body weight, body weight gain, food consumption, food utilization, hematology parameters (hemoglobin concentration, erythrocyte count, total and differential leukocyte counts), urinalysis parameters (reducing substances, albumin, microscopic elements), gross pathology, organ weights (liver, kidneys, spleen, gonads, heart, brain, thyroid gland, adrenal gland), tumor incidence (organs examined not specified), and microscopic pathology (heart, lung, trachea, liver, pancreas, stomach, small intestine, colon, spleen, lymph node, kidney, urinary bladder, testis, ovary, prostate, uterus, pituitary, adrenal gland, salivary gland, thyroid gland, parathyroid gland, skeletal muscle, bone marrow, brain).
	No increase in mortality occurred and the only clinical signs were generalized inactivity and weakness in animals dying on study. A significant decrease in body weight gain was noted in the 1% dose group at the interim sacrifice (12 months) only. Body weights were also decreased in this

	group at the 12-month time point. After 24 months, no effect of treatment on body weight or body weight gain was observed. Food consumption was decreased in the high-dose group, but food utilization was unaffected. It was suggested that the body weight and food consumption effects were related to the palatability of the test diet. No effects on hematology, urinalysis, organ weights, and gross and microscopic pathology were reported. Reproductive organs ( <i>i.e.</i> , testes, ovaries, uterus) showed no evidence of toxicity at any dose level. The tumor incidence and tumor types were similar in the test and control groups. Based on these data, it appears that the NOEL was 0.2% (approximately 200 mg/kg/day).
Data Quality	Valid without restriction – Klimisch Code 1b
References	Kay, J.H. 1962. Two-year chronic oral toxicity of [trade name deleted] hydrogenated rosin – albino rats. Industrial Bio-Test Laboratories, Inc., Northbrook, Illinois.
	World Health Organization (WHO). 1990. Principles for the Toxicological Assessment of Pesticide Residues in Food.